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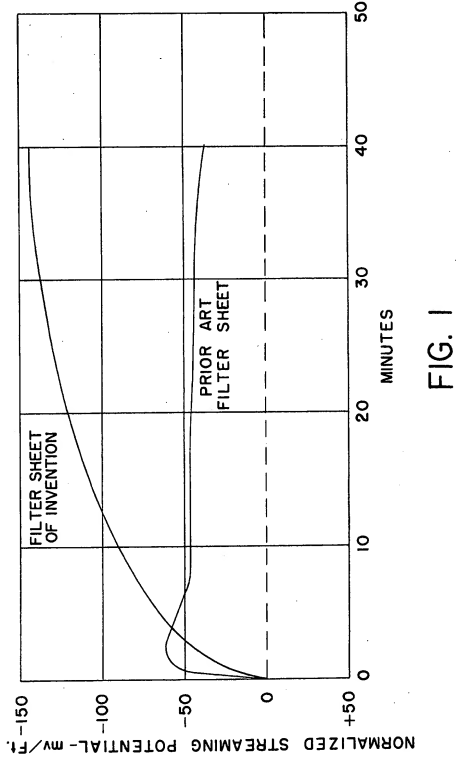
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## (54) Filter and method of making same

(57) A filter sheet comprises fine inorganic particles and a self-bonding matrix of cellulose fiber, the surfaces of at least one of which are modified by colloidal silica made cationic by the deposition on it of an oxide or hydroxide of a polyvalent metal, particularly aluminium. The sheet is made by deposition from a slurry in water, and the cationic modification may be before or after sheet formation.



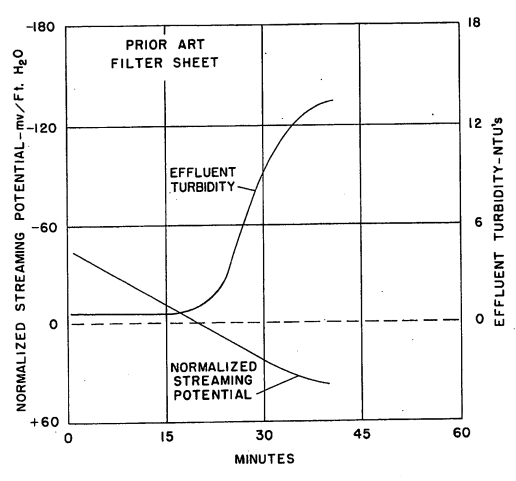


FIG. 2

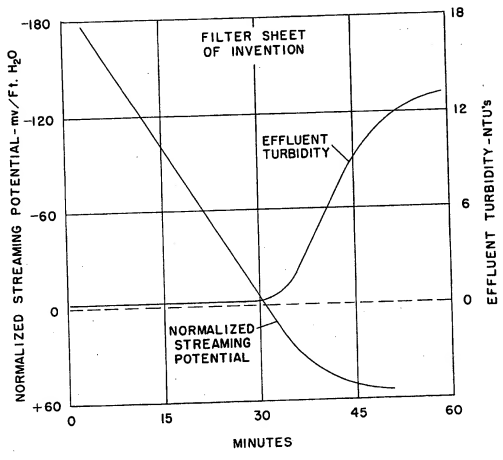


FIG. 3

## **SPECIFICATION**

## Filter and method of making same

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5	This invention relates to filtration and, more particularly, to the removal of submicron contaminants from aqueous systems, utilizing filter media sheet comprising high levels of particulate filter aids.	5
10	The filtration of fine particle size contaminants from fluids has been accomplished by the use of various porous filter media through which the contaminanted fluid is passed. To function as a filter, the media must allow the fluid, commonly water, through, while holding back the particulate contaminant. This holding back of the contaminant is accomplished by virtue of the operation, within the porous media, of one or both of two distinctly different filtration mechanisms, namely (1) mechanical straining and (2) electrokinetic particle capture. In mechanical straining, a particle is removed by physical entrapment when it attempts to pass	10
	through a pore smaller than itself. In the case of the electrokinetic capture mechanisms, the particle collides with a surface face within the porous filter media and is retained on the surface by short range attractive forces.	15
	With the exception of microporous polymeric membranes, the porous filter media known to the art as being suitable for the filtration of fine particle size contaminants are comprised of fiber-fiber or fiber-particulate mixtures formed dynamically into sheet by vacuum felting from an aqueous slurry and then subsequently drying the finished sheet. In those fibrous filter media that depend upon mechanical straining	•
20	to hold back particulate contaminants, it is necessary that the pore size of the filter medium be smaller than the particle size of the contaminant to be removed from the fluid. For removal of fine, submicronic contaminant particles by mechanical straining, the filter media need have correspondingly fine pores. Since the pore size of such a sheet is determined predominantly by the size and morphology of the materials used	20
25	to form the sheet, it is necessary that one or more of the component materials be of a very small size, such as small diameter fibers. See, for example, any of Pall U. S. Patents 3,158,532; 3,238,056; 3,246,767; 3,353,682 or 3,573,158.	25
20	As the size of the contaminants sought to be removed by filtration decreases, especially into the submicron range, the difficulty and expense of providing suitably dimensioned fiber structures for optimum filtration by mechanical straining increases. Accordingly, there is considerable interest in the use of fine particulates such as diatomaceous earth.	30
30	However, for such materials it is necessary to provide a matrix in order to present a coherent handleable structure for commerce and industry. Thus, at least one of the component materials in the sheet is a long, self-bonding structural fiber, to give the sheet sufficient structural integrity in both the wet "as formed" and	
35	in the final dried condition, to allow handling during processing and suitability for the intended end use.  Unrefined cellulose fibers such as wood pulp, cotton, cellulose acetate or rayon are commonly used. These fibers are typically relatively large, with commercially available diameters in the range of six to sixty micrometers. Wood pulp, most often used because of its low relative cost, has fiber diameters ranging from	35
40	fifteen to twenty-five micrometers, and fiber lengths of about 0.85 to about 6.5 mm.  Filter media sheets are conveniently formed by vacuum felting from an aqueous slurry of the component material. The vacuum felting is performed on a foraminous surface, normally a woven wire mesh which, in practice, may vary from 50 mesh to 200 mesh, with mesh openings ranging from 280 micrometers to 70 micrometers respectively. Finer meshes are unsuitable because of clogging problems and/or structural	40
45	The size of the openings in the foraminous vacuum felting surface, and the pore size of the cellulose fiber matrix of the formed sheet, are quite large in comparison to some or all of the dimensions of the fine fiber or particulate components required to produce the desired submicronic filter media sheet. Retention of such fine components during the vacuum formation of the filter media sheet is difficult, and imposes severe	45
50	constraints on the choice of such materials, the specific details of the process utilized to form the filter media sheet, and, most important, upon the level of filtration performance that may be attained. Fine fibers, whose length may be large in comparison to their diameter, present less of a problem and tend to be retained reasonably well. Fine particulates, on the other hand, tend to show very poor rentention during sheet	50
55	formation.  Flocculation with polymeric retention aids, or coagulation has been used as a means of improving retention of fine particulates, in effecting the grouping of particles to offer an effective large dimension. However, filter sheet prepared from a well-flocculated slurry will have a broad particle size distribution, with small pores occurring inside the flocs, and large pores occurring between the flocs. The existence of these larger pores will limit the ability of the filter media sheet to remove fine contaminants. The use of flocculation	55
60	to achieve high retention in filter media is therefore somewhat counter-productive. It is, of course, possible to apply hydrodynamic shear forces, breaking up the flocs, and further charge modify until the system assumes a stable disperse form. This does achieve a relatively uniform sheet of narrow pore size distribution. However, the retention of the particulates in such a system is very low, leading	60
65	to concomitant reduction of filtration efficiency. In addition to controlling the dispersion characteristics (and therefore the porosity of the sheet) and providing wet strength, charge modifiers are employed to control the zeta potential of the sheet constituents and maximize performance in the electrokinetic capture of small charged contaminants. In practice, cationic	65

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charge modifiers are employed since most naturally occurring contaminant surfaces are anionic at fluid pH of practical interest. Thus, a melamine-formaldehyde cationic colloid is disclosed for filter sheets in U. S. Patents 4,007,113 and 4,007,114.

The use of such filter systems with biological fluids poses special problems, among them the possibility of introducing to the fluid impurities resulting from loss of or a breakdown in filter elements. While certain levels of particular impurities may be tolerable in some systems, organic extractables pose especially sensitive problems in the filtration of foods and pharmaceutical products. In filter systems composed of cellulose fiber as a matrix for particulate filter aids modified with an organic cationic resin, organic extractables are naturally primarily traceable to the resin. Selection of the charge modifying resin can alleviate the problem, even under relatively stringent conditions of use including sanitization and sterilizable procedures. Even in the absence of meaningful levels of extractables, however, many resins of choice are subject to discoloration in use, tending to limit their marketability for food and drugs.

Further, even low levels of certain organic extractables are unacceptable in some systems, and

Further, even low levels of certain organic extractables are unacceptable in some systems, and accordingly it is desirable for this reason and that of aesthetics to wholly remove the organic charge modifier resin from the filter construction. At the same time, it is necessary for the removal of submicron negatively charged contaminants to retain the positive charge potential afforded by a charge modifying resin.

It is accordingly an object of the present invention to provide charge modified filter media sheets of enhanced filtration performance, especially for the removal of submicron contaminants from aqueous systems at high efficiency.

Another object is to provide charge modified filter media characterized by low organic extractables over a wide range of filtration conditions.

A still further object is the provision of filter media effective across the spectrum of biological liquids and, particularly, ingestables such as foods and drugs.

These and other objects are achieved in the practice of the present invention as described hereinafter.

In accordance with the present invention, charge modified filter media sheets are prepared utilizing an inorganic charge modifier constituted by cationic colloidal silica. The filter media sheet, preferably formed by vacuum felting of a cationically disperse aqueous slurry comprising beaten cellulose fibers and fine particulate shows a uniform, high porosity, and fine pore size structure with excellent filtration and flow characteristics. Thus, the sheets comprise cellulose fiber as a matrix, and particulate filter aid, the surfaces of at least one of which have been modified with cationic colloidal silica. Those sheets are free of extractables, such as formaldehyde or amines originating with organic resinous charge modifiers, and are free of discoloration, such that the sheets are useable under any sterilizing conditions and may be employed safely

and effectively with potables or ingestables such as food or drugs.

The cationic colloidal silica is an aqueous dispersion of positively charged colloidal particles consisting of a dense silica core coated with a positively charged polyvalent metal-oxygen compound typically stabilized with a counterion. Illustrative such materials are disclosed in U. S. Patent 3,007,878 incorporated herein by reference. Other surface modified cationic colloidal silica materials are known for insolubilization of enzymes, for example as shown in U. S. 3,796,634 but are characterized by the presence of organic modifying resins and hence are contraindicated for use in conjunction with the present invention.

While the principal requirement for this component of the filter system is that it functions as a charge modifier and dispersion agent, it should also be able to interact with the matrix to provide strong bonding as by cross-linking. U. S. Patents 3,775,141 and 3,785,838 disclose refractory compositions comprising inorganic fibers bonded with cationic colloidal silica into integral sturctures which may also comprise a proportion of inorganic filler such as perlite. It is surprising then, in that the bonding of organic to inorganic materials is more difficult to achieve, to find that strong bonds are achieved with cellulose fiber upon curing with cationic colloid silica. However, the bonding once achieved is relatively more inert to its surrounding environment, and loss of the material by hydrolytic or solvolytic action, believed to be among the causes of impurities or extractables generated in other systems, is obviated.

In preferred embodiments of the invention, relatively high loadings of fine particulates such as diatomaceous earth or perlite, to 50-70 percent of more by weight of the sheet, are employed. Without wishing to be bound by an essentially hypothetical elucidation, it is believed that surface modification of these materials with cationic silica colloid, particularly at these high loadings, contributes to the integrity of the overall structure, and may be attributable to the formation of some siliceous, or inorganic interbondings, interengaging the relatively low level (10-20%) of cellulose fibers comprising the total sheet weight in such embodiments, with the particulates by way of the cross-linking action of the active hydroxyl sites provided by the colloidal silica.

The present invention is further illustrated by the accompanying drawings, in which:

Figure 1 is a graph of normalized streaming potential vs. time, comparing equilibrium flush out curves for a prior art filter sheet, and a filter sheet manufactured in accordance with the invention.

Figures 2 and 3 are graphs of normalized streaming potential and effluent turbidity vs. time, comparing monodisperse latex contaminant challenge tests for a prior art filter sheet, and a filter sheet manufactured in accordance with the invention.

The filter media sheets of the invention are prepared from cationically modified filter elements, usually in the form of a cationically disperse aqeuous slurry comprising cellulose fiber and optimized levels of fine particulate such as diatomaceous earth or perlite. The filter elements may be cationically modified in the

slurry and the sheet prepared dynamically by vacuum felting, and drying, or the filter elements may be pretreated and formed into sheet media. A special feature of the invention is the provision of filter media sheet in which the level of particulate retained is enhanced as compared to sheet prepared conventionally.

The state of refinement of a wood pulp fiber is determined by means of a "freeness" test in which

5 measurement as the flow rate through a forming pad of the fibers on a standard screen is determined, most commonly utilizing the "Canadian Standard Freeness Tester". In this method, the quantity which is measured is the volume of water (expressed in ml.) which overflows from a receiver containing an orifice outlet at the bottom. The Canadian Standard Freeness measurements are employed in the present specification. Coarse unbeaten wood pulp fibers produce high drainage rates into the receiver from the screen resulting in large overflow volumes, and hence record a high freeness. Typical wood pulps show Canadian Standard Freeness values ranging from +400 ml. to +800 ml. In paper or filter media manufacture, such pulps may be subject to mechanical refining processes such as beating, which tends to cut and/or

fibrillate the cellulose fibers. Such beaten fibers exhibit slower drainage rates, and, therefore, lower freeness. In accordance with the present invention, such beaten pulp is preferably employed in the self-bonding matrix for the filter media. The Canadian Standard Freeness of the pulp system will vary with pulp selection, and may be reflective of varying states of subdivision or refinement, as where different pulps or differently beaten pulps are combined for sheet formation, but the beaten pulp will be employed to provide a composite or average value ordinarily ranging from 100 to 600 ml., with lower values e.g., 200-300 ml.or less being

preferred for higher solids retention.

The wood pulp may comprise as little as 10 percent by weight with up to 20 to 30 percent, by weight of the total, being preferred to provide filter media sheet with structural characteristics suitable for industrial filtration applications.

Performance is enhanced by maximizing the amount of fine particulate in the filter media sheet. While as little as 10 percent of a fine particulate will result in noticeable improvement in filtration performance of either type of media, optimum performance is achieved by utilizing the maximum amount of fine particulate. For industrial filtration, structural characteristics suggest a practicable maximum of about 70 percent by weight. Of course, for less demanding applications, somewhat higher levels will be possible. Generally,

levels of 50-70 percent by weight are employed.

There are various types of fine anionic particulates that are suitable for the intended purpose, including diatomaceous earth, perlite, talc, silica gel, activated carbon, molecular sieves, clay, etc. Functionally, the fine particulate should have a specific surface area in excess of one square meter/gram and/or particle diameters of less than 10 microns. In a broad sense, any fine particulate may be suitable (such as J. M. Filter Cel, Standard Super Cel, Celite 512, Hydro Super Cel, Speed Plus and Speedflow; Dicalite 215 and Dicalite 416 and Dicalite 436) and may be evaluated by techniques well-known to the art. Siliceous materials are preferred, and from the stand-point of size, morphology, cost, fluid compatibility and general performance characteristics, the finer grades of diatomaceous earth and perlite filter aids exhibiting a mean particle size of less than 5 microns are most preferred. In many cases, mixtures of more than one type of fine particulate such as diatomaceous earth/perlite for example, in proportion by weight of from about 80/20 to 20/80 give

better filtration performance or better cost/performance characteristics than that achieved by the use of any single type by itself. Similarly, mixtures in all proporations of relatively coarse and fine particulates, e.g., 50/50 parts by weight of 10 and 5 micron diameter particulates may be used.

Suitable cationic colloidal silica materials include those of U. S. Patent 3,007,878, 3,252, 917, 3,620,978, 3,719,607 and 3,956,171 all incorporated by reference. These are aqueous dispersions of positively charged colloidal particles consisting of a dense silica core coated with a positively charged polyvalent metal-oxygen compound, e.g., of the class consisting of metal oxides, metal hydroxides and hydrated metal oxides of metals having a valence of 3 or 4, preferably aluminum and titanium. Most preferably, the dispersion is acidic, and the coating is of polymeric alumina species. Typically, the mole ratio of aluminum to silica on the surface is about 1:1, and the dispersion (which has been commercially available as Ludox Positive Sol 130M, from E. I. duPont de Nemours & Co.) is stabilized with a counterion, as described in the aforesaid U. S. Patent 3,007,878. The dispersion has been supplied at 30% solids, stabilized with chloride ion (1.4%, as NaCl) for use in the pH range 3.5 to 5.5.

The colloidal particles exhibit a surface area of about 150-225 m<sup>2</sup>/g by nitrogen absorption, a particle diameter of about 15-16 mu, and a molecular weight of about 5 to 18 million by light scattering.

In its preferred form, the characteristics of the silica aquasol are further modified to higher levels of polymeric alumina species calculated as alumina from 13 to 15% or more, in the stable range based upon the colloidal solids. The coating, or overcoating, may be achieved by simply treating with an appropriate aluminum compound, e.g., basic aluminum chloride, as described in U. S. 3,007,878, or another source of polybasic aluminum cations. The alumina in such systems exists as a surface coating and, to the extent it exceeds the available surface area, as free alumina in solution. The free alumina may, of course, also serve as a coating for virgin filter elements, e.g., particulate present and systems so prepared offer improved resistance to autoclaving and hot water flushing conditions together with added wet strength. Where desired, the resulting colloidal dispersion may be, and customarily is treated to remove excessive electrolyte, as by dialysis, in order to achieve storage stability. Filter sheets prepared with the preferred

cationic colloidal silica exhibit uniformly acceptable sterilization stability under stringent conditions, e.g., 65 autoclaving at 15 psi, 121°C. for 1 hour.

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5	In paper production, where cationic charge modifiers are sometimes used, the objective is reduction of charge to approximately the isoelectric point to maximize efficiency in interfelting of fiber. For filtration, maximum charge is desired to enhance removal of charged particles by electrokinetic mechanisms. In the present case the surface charge of at least one of the negatively charged filter elements, i.e., cellulose and particulate is reduced to render the surface less electronegative and optionally (and preferably) reversed by the deposition of sufficient cationic charge modifier to render the surface electropositive, to provide at least certain electropositive regions or sites within the filter sheet. In order to accomplish charge reversal, of course, one proceeds through the isoelectric point, and then positive charge buildup is accomplished to the	5	
10	maximum practical level.  The amount of charge modifier employed in the present invention is thus preferably that sufficient to at least provide a cationically disperse system, i.e., a system in which no visible flocculation occurs at ambient conditions in the absence of applied hydrodynamic shear forces. The system therefore comprises essentially	10	
15	discrete fiber/particulate elements exhibiting a positive charge or zeta potential relatively uniformly or homogeneously distributed in and throughout the aqeuous medium. The specific level will, of course, vary with the system and the modifier selected but will be readily determined by one skilled in the art. For example, the inflection point on a plot of particulate retention vs. amount of charge modifier approximates	15	
	the minimum level for better performance. Thus a 5-6 percent level is appropriate for cationic colloidal silica applied during sheet formation, although lower amounts, e.g., 3-4 percent may be sufficient where the filter elements are precoated. Although additional modifier may be employed to advantage where desired, this		
20	level represents the best balance on a cost/performance basis. Premodified filter elements e.g., cellulose fiber and particulate precoated with charge modifier may of course be incorporated in any manner into filter sheets with similar results, and where a cationically disperse slurry is not employed, charge modification will be commensurately reduced by control of modifier levels.	20	
25	The charge modification effected is demonstrable in measurements of surface zeta potential, and in improved filtration efficiency for negatively charged particles in liquid systems.  The slurry of pulp and particulates is formed in any suitable manner. The sequence of adding these components to water to form the initial slurry appears to be relatively unimportant. The consistency of the	25	
30	slurry will represent the highest possible for a practical suspension of the components, usually about 4 percent. The system is subjected to hydrodynamic shear forces as by a bladed mixer, and the charge modifier is then added to the slurry.  The shear level is not critical i.e., any otherwise suitable shear rate or shear stress may be employed	30	
	having regard for available equipment, preferred processing times etc. but is selected and employed simply to break up the flocs and maintain the system in a dispersed condition during treatment. Of course, upon the formation of a cationically disperse slurry, the system is free of floc formation even in the absence of applied		
35	5 shear.  After charge modification, the slurry is diluted with additional water to the proper consistency required for vacuum felting sheet formation, ordinarily 0.5 to 2 1/2 percent, depending upon the type of equipment used to form the sheet in a manner known to the artisan. The slurry is formed into a sheet and oven dried in	35	
40	standard manner. The performance of the sheet is related to the drying parameters and optimized conditions of may reflect energy considerations or desired thermal history consistent with minimization of unnecessary exposure to elevated temperatures, especially as the decomposition or scorch point for the system is	40	
4!	approached. In accordance with a preferred embodiment of the invention, filter media sheets are formed from filter elements, i.e., particulate and a self-bonding matrix of cellulose pulp at least one of which is charge modified, the pump being a system incorporating beaten pulp to provide a Canadian Standard Freeness of up to 600 ml., preferably less than 300 ml. e.g., 100-200 ml. the charge modifier consisting of cationic silica comprising at least 20% surface alumina and being applied in a proportion to reduce electronegativity of the	45	
5	surface, preferably to achieve charge reversal beyond the isoelectric point, e.g., to an add-on level of about 6% by weight. Filter media sheets so prepared may be autoclaved, hot water flushed or otherwise treated at 0 elevated temperature to sanitize or sterilize the structure. In addition to having special utility for the filtration of biological liquids, these sheets also may be employed for filtration without flush out delay, as ions present in the structure have been removed during the sanitization or sterilization procedure.	50	
5	The performance benefits of the cationic colloidal silica may be seen dramatically illustrated in the drawings, representing the results of testing conducted as described in Example V. The cationic colloid silica of the invention retains hi charge and contaminant removal capacity and has longer effective life than the melamine-formaldehyde cationic colloid of U. S. Patent 4,007,113.  The filter media sheets may be subjected to standardized testing reflecting performance in use,	55	•
6	represented herein by the following:	60	
9	In this test, contaminated fluid of a specified turbidity is drawn through the test filter media by a constant		

In this test, contaminated fluid of a specified turbidity is drawn through the test filter media by a constant vacuum under standard conditions, effluent turbidity measured (using a Hach model 2100A Turbidimeter)

and expressed as percent filtration efficiency when compared to the inlet turbidity, calculated as

Efficiency = Turbidity In -.

Turbidity Out

Turbidity In

Test contaminant is Hyplar (produced by Grumbacher) a polydisperse acrylic latex produced by emulsion polymerization and comprising colloidal polymer particles ranging from 0.05 to 1.0 micron. Contaminant level is 25-200 NTU (Nephalometric Turbidity Units) in distilled water, at pH 6.5-7.0. Filter sheet is cut into 57 mm dia. discs, and then placed in a Millipore 47 mm vacuum filter holder. 100 ml. of the prepared contaminant dispersion is then filtered through the disc using a 23 in. Hg vacuum.

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#### 10 Membrane Protection Test

In this test, contaminated fluid is pumped under standard conditions through test filter media and a membrane in series at a constant flow rate, and differential pressure with time recorded. The time or total volume of flow passed at a defined pressure increase is a measure of the life of the prefilter, and interrelates satisfactorily with preformance in use. Typically, a 47 mm. 0.22 micron membrane is employed at a flow rate of 225 ml./min. Test contaminant is the same Hyplar polydisperse acrylic latex referred to above.

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of 225 ml./min. Test contaminant is the same Hypiar polydisperse acrylic latex referred to above. Contaminant level is 5-50 NTU (Hach Turbidimeter, Model 2100A). The test is continued until the differential pressure across either the membrane or the test filter paid exceeds 5-10 psid. Membrane protection times of less than a few minutes indicates no practically useful effect.

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#### 20 Oil Flow Test

As a measure of the porosity of the filter media sheets, 100 ssu oil is pumped through the sample sheet until a differential pressure drop of 5 psid is attained, at which point the flow rate (ml./min.) is recorded.

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#### Normalized Streaming Potential

25 The measurement of streaming potential is a conventional means of determining zeta potential i.e., the electric potential excess of the surface, and the surrounding fluid to the hydrodynamic shear plane, over the bulk potential of the fluid. In the present test, streaming potential values are determined, and normalized for differing pressure drop in the media being tested, expressing the results in units of millivolts per foot of water. The filter media is evaluated by flushing out the filter media with water until the measured streaming potential achieves a relatively stable maximum value. At this point, the filter media has ceased to contribute any significant ionic species to the water, i.e., the inlet resistivity equals the outlet resistivity.

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The filter media test cell is based on the design of Oulman, et al. JAWWA 56:915 (1964). It is constructed from Lucite having an effective area of 3.14 square inches (2" diameter) and is equipped with platinum black electrodes. Water and mercury manometers are used to measure the pressure drop across the media being evaluated. Streaming potential values (by convention, of opposite sign to the zeta potential and the surface charge) are measured with a high impedance volt meter. The influent and effluent resistance are monitored with conductivity flow cells (cell constant = 0.02/cm) using a resistance bridge.

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Upon the attainment of equilibrium streaming potential (i.e., after flush out) contaminant challenge tests may be performed in the same system, using an aqueous dispersion of a single size monodisperse latex, while measuring the 90° light scattering intensity (selected for high relative sensitivity for particle diameters in the 0.1 to 1.0 micrometer range) of the influent and effluent streams, providing a quantitative determination of the filtration efficiency of the filter media. Effluent turbidity is measured using a Moniteck

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Turbidity Meter. Inlet turbidity is selected to range from 15 to 20 NTU, using a Dow Diagnostics 0.109 micrometer emulsion polymerized polystyrene latex dispersion, and flow rate is maintained relatively constant. The tests above are described in more detail in a paper presented at the 71st Annual AICHE meeting (1978): "Measuring the Electrokinetic Properties of Charged Filter Media", Knight et al.

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In the following examples which further illustrate this invention; proportions are by weight, based upon total pulp and particulate, excluding charge modifier.

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#### 50 Example I

A series of filter sheets were prepared utilizing Weyerhauser Coho Kraft pulp, beaten to the levels indicated below, and Grefco Dicalite 416 Perlite, having a mean particle size of 3.9 microns.

The charge modifier employed in these runs was a cationic colloidal silica (30% solids) containing about 15% alumina, based upon the weight of colloidal solids (determined by atomic absorption at 309 nanometer 55 with a hollow cathode lamp and nitrons oxide-acetylene flame), having a surface area of about 220 m²/g (nitrogen adsorption) a particle diameter of about 15-16 mu and a molecular weight of about 5-18 million (light scattering).

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The total input weight (bone dry basis) of the component materials was 80 grams, exclusive of charge modifier. A constant proportion of pulp (30 percent by weight, or 24 grams) and particulate (70 percent by 60 weight, or 56 grams) was maintained. The components were added to water in a 1 liter polyethylene bucket, with strong agitation, to form an aqueous slurry at two percent consistency, and the charge modifier added. (The system was subjected to hydrodynamic shear by action of Hei-Dolph stirrer (Polyscience Inc.), having 4 propeller blades, rotating at about 700 rpm on setting 2.) The slurry was subsequently diluted to 0.5 percent consistency and vacuum felted into a sheet ranging from about .160 to .200 inch thickness (depending upon 65 retention) in a nine inch by twelve inch hand sheet apparatus utilizing a 100 mesh screen. The sheet was

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subsequently removed, dried in a static oven at 350°F, until constant weight was achieved, and the final weight recorded. Comparison of the final sheet weight with the total input material weight allowed determination of total solids retention in the sheet. Filter sheets were subjected to filtration (membrane protection) and oil flow tests, as described above. Results are set forth in Table I.

TABLE I

10	Sheet No.	Pulp Freeness (CSF)	Charge Modifier Content (Wt.5	Oil Flow %)(Ml/Min.)	Minit.△P (PSID)	embrane protec Time (Min.)	etion test ( 25 PAD∆∆P (PSID)	NTU) MEMB.∆∆P (PSID)	10
	1	660	6	-	1.1	127	0.6	5.0	
15	2	660	6	•	0.9	123	1.0	5.0	15
	3	660	0	135	0.8	14	1.2	5.0	
	4	500	6	-	1.5	110	10.0	0.9	20
20	5	500	6	-	1.2	115	10.0	0.5	20
	6	520	0	67	1.0	0	1.5	5.0	
25	7	350	6	-	1.5	73	10.0	0.8	25
	8	350	6	-	1.9	. 72	10.0	0.3	
	9	400	0.	111	0.3	<b>5</b> .	1.0	5.0	30
30 <u>.</u>	10	258	6	-	1.6	50	10.0	0.4	
	11	258	6	•	1.8	69	10.0	0.4	
35	12	320	0	98	1.0	4.0	1.0	5.0	35
	13	167	6	-	2.1	48	10.0	0.5	
	14	167	6	-	2.2	70	10.0	0.5	40
40	15	200	0	58	1.0	1.0	1.0	5.0	40
	16	100	6	-	2.8	35	10.0	0.2	
45	17	100	6	-	2.8	48	10.0	0.3	45
	18	110	. 0		N	IOT TESTED -		<b></b> →	

As will be seen from the above reported testing, samples which were not charge modified evidenced essentially no membrane protection. Failure in the systems of lowest freeness was by pad plugging, with little increase in membrane pressure drop.

Runs were conducted in the manner of Example I, employing as the particulate Grefco Perlite 426, having a mean particle size of 4.2 microns, and 436 having a mean particle size of 6.0 microns and a constant (6%) level of Wesol PA, a cationic colloidal silica charge modifier available from Wesolite Corp., Wilmington Delaware (4.0% alumina, 22.5% silica, 30% solids). Results are set forth in Table II.

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				Membrane Protection Test (50 NTU)				
	Sheet No.	Pulp Freeness (CSF)	Charge Modifier Content (Wt. %)	Init.△P (PSID)	Time (Min.)	△△PAD P (PSID)	MEMB.△△P (PSID)	5
•	19 (436 Perlite)	185	6	0.7	17.1	5.0	3.0	10
	20 (426 Perlite)	130	6	1.4	20.6	5.0	0.5	

15 Example III

A. Particulate filter aid was precoated in an aqueous slurry at a 2.5% consistency with the below indicated levels of Wesol PA cationic colloidal silica charge modifier over a contact time of about 15 minutes, isolated, drained and dried at 250°F. for 30 minutes.

The treated particulate filter material was slurried in 100 ml. of water and filtered through a porous fritted 20 glass holder base in a Millipore 47 mm, vacuum filter holder, until a 1/4 inch thick cake was formed, and

filtration efficiency tests were then performed. Results are set forth in Table III.

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5	Sample No.	Particulate Type	Charge Modifier Content (Wt. 5	Inlet Turbidity (NTU) %)	Efficiency Test Effluent Turbidity (NTU)	Efficiency (%)	5
	20	DE 215 <sup>1</sup>	0	200	50	75.0	٠
10	21	DE 215	1 .	200	15	92.5	10
	22	DE 215	3	200	1.7	99.2	
	23	DE 215	6	200	3.8	98.1	15
15	24	PERLITE 416	0	200	100	50.0	
	25	PERLITE 416	1	200	69	65.5	
20	26	PERLITE 416	3	200	0.96	99.5	20
	27	PERLITE 416	6	200	5.0	97.5	
	28	PERLITE 426	0	200	100	50.0	25
25	29	PERLITE 426	1	200	1.4	99.3	
	30	PERLITE 426	3	200	1.7	99.2	
30	31	PERLITE 426	6	200	8.0	96.0	30
	32	PERLITE 436	0	100	78.0	25.0	
	33	PERLITE 436	1	100	1.4	98.6	35
35	34	PERLITE 4106 <sup>2</sup>	0	50	45	10.0	•
	35	PERLITE 4106	1	50	8.0	98.4	
40	36	PERLITE 4106	3	50	1.1	97.8	40
	37	PERLITE 4106	6	50	3.5	93.0	
	<sup>1</sup> DE 215 is G	refco Dicalite calcine	d diatomaceo	us earth, having a m	ean particle size of 2	.7 microns.	45
45							75

<sup>&</sup>lt;sup>2</sup>Perlite 4106 is Grefco Dicalite perlite having a means particle size of 10 microns.

The improvement in filtration performance with charge modification is dramatic, especially with the larger particulates.

Example III

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B. An aqueous slurry of the precoated particulate prepared as set forth above is alternatively formed into a filter sheet comprising 30 percent by weight beaten pulp as a matrix, and dried. The pulp system employed exhibited a freeness of 130 C.S.F. Membrane protection and oil flow tests were performed, with results set 55 forth in Table IV.

#### TABLE IV

					ion Test (25 NTU)					
•	5	Sheet No.	Particu- late Type	Charge Modifier Pretreat (Wt. %)	Oil Flow (Ml/Min.)	Init. △P (PSID)	Time (Min.)	PAD∆∆P (PSID)	MEMB. △△P (PSID)	5
		38	416 PERLITE	0	21	2.0	0 .	0	10	
	10	39	416 PERLITE	6	32	1.7	9.0	0.3	5	10
		40	4106 PERLITE	0	262	0.6	0	0	10	
	15	41	4106 PERLITE	6	266	0.6	8.3	0	<b>5</b> ·	15

C. A preformed sheet comprising a 130 C.S.F. pulp system (30% by weight) and Perlite 416 particulate was impregnated with a 6% by weight solution of Wesol PA cationic colloidal silica charge modifier, dried 20' and cured. In filtration tests, membrane protection time was 5.0 minutes, and failure was by pad plugging at 5 psid.

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Example IV

A. In the manner of Example III A, mechanically defibered cellulose (Solka floc) was precoated with 25 Wesol PA cationic colloidal silica charge modifier, dried and cured, formed into a filter cake, and tested for filtration efficiency. The results are set forth in Table V, as follows.

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## TABLE V

		Sample No.	Particulate Type	Modifier	Inlet Turbidity (NTU)	Efficiency Test Effluent Turbidity (NTU)	Efficiency (%)	30
	35	42	Solka-Floc BW-20	0	10	6.9	21.0	35
		43	Solka-Floc BW-20	1 .	10	7.9	31.0	40
	40	44	Solka-Floc BW-20	3	10	1.5	85.0	40
	45	45	Solka-Floc BW-20	6	10	2.5	75.0	45
		46	Solka-Floc BW-200	0	10	5.9	41.0	
•	50	47	Solka-Floc BW-200	3	10	2.2	78.0	50

B. A filter sheet was formed from a slurry consisting of 30% untreated cellulose pulp and 70% pretreated defibered cellulose from Example IV A, and tested as described in Example III B, with the results set forth in Table Vi, as follows.

<b>T</b> A			1/1
IΑ	В	Ŀ	VI

					•			
5	Sample No.	Particulate Type	Charge Modifier Pretreat (wt %)	Membrane P Init. PAD △P (PSID)	rotection (5NT Time · (Min.)	TU) PAD∆∆P (PSID)	`Memb. ∆∆P (PSID)	5 ·
10	48	Solka-Floc BW-20	0	0.5	1.75	0	10	: 10
10	49	Solka-Floc BW-20	1	0.3	1.75	0	5	
15	50	Solka-Floc BW-20	3	0.3	3.00	0	10 .	15
	51	Solka-Floc BW-20	6	0.3	10.00	0	5	
20					•	•		20
	modified t	xample, the perform	nance of a prior art n ee U. S. Patent 4,007 Vesol PA) in accorda	,113) was com	ipared to filter	onic colloid (Pa media sheet cl	arez 607) charge harge modified	
25	A. Filte and 30% t manner b optimized	er media sheets wer by weight of a 50/50 y preparing a cation I charge modifer lev	vesor PA) in accorda e prepared containi admixture of diatom ically disperse aque els (7% for Parez 607	ng 70% by wei naceous earth ous slurry, vai	ght of a cellulo and perlite and cuum felting ar	nd oven drying	, except that	25
30	employed.  30 Employing test conditions detailed above, normalized streaming potential values were determined over time, and equilibrium flush out curves plotted for the respective filter media, compared in Figure 1. As will be seen, the prior art (melamine-formaldehyde) media exhibits an anomolous flush-out curve, peaking out very rapidly and then slowly decaying with time. The media sheet prepared in accordance with the invention							
35	exhibited an increasingly negative normalized streaming potential which stabilized at a high equilibrium  stabilized at a high equilibrium value indicative of a high positive surface charge.  B. The same filter media were then challenged with contaminated liquid (0.109 micron Dow Diagnostics latex dispersion; 5.5 pH), and filtration efficiency and normalized streaming potential plotted against time in Figures 2 and 3 comparing respectively the prior art (melamine-formaldehyde cationic colloid) system							35
40	(Figure 2) latex part normalize	with the cationic co icles are, initially, es ad streaming potent	illoidal silica modifie sentially quantitativ ial decays linearly fr	rely removed b rom a negative	by electrokinet value, throug iches and pass	ic capture and h zero, and the es through zer	adsorption. The n asymptotically o, the effluent	40
4!	turbidity approach The filt	starts to decay (brea les the inlet turbidity er media sheet of th	kthrough). This incr indicating that all o e invention (Figure 3)	ease continue of the latex is p 3) retained its f	s until the enit assing through litration perfo	n the filter med rmance signific	ia. cantly longer	45
5	'Biological liquids' as that term is employed in the specification and claims, is a liquid system which is derived from or amenable to use with living organisms, and ordinarily handled and processed under sanitary or sterile conditions, therefore requiring sanitized or sterilized media for filtration. Included are isotonic solutions for i.m. or i.v. administration, solutions designed for administration per os, as well as solutions for topical use, biological wastes or other body fluids which may comprise filterable bodies such as impurities and hacterial viruses or pyrogens which are desirably isolated or separated for examination or							50,
5	disposal by immobilization or fixation upon or entrapment within filter media. It is filter media sheets in accordance with the invention may be employed alone or in combination with other such media to treat pharmaceuticals such as antibiotics, saline solutions, dextrose solutions, vaccines, blood plasma, serums, sterile water or eye washes; beverages, such as cordials, gin, vodka, beer, scotch, whisky, sweet and dry wines, champagne or brandy; cosmetics such as mouthwash, perfume, shampoo, hair tonic, free grown or shaving lettin; food products such as vinegar, vegetable oils, extracts, syrups, fruit juices,							55
6	electropl retention The filt solutions	lating solutions, clean of submicronic par ter media of this inve s and especially the	ls; chemicals such a ning compounds, s ticles, removal of ba ention are particular removal of the most	olvent purifica acterial contan Iv suitable for	ition and lubric ninants and res the removal o	cating oils; and solution of coll f pyrogens froi	oidal hazes. m biological	60
6	parenter 5 Filter r	al solutions. nedia of the inventio	on were evaluated in	this regard u	sing two test p	rocedures.		65

5	The first involved the passage of small volumes of test solution (10-50 ml.) through 0.9 cm² disc filters. The test solution was passed through the filters with the aid of a syringe at a constant flow rate. In large volume experiments usually ten liters of test solution was passed through 3.9 cm² filter at a flow rate of 3.05 gal./sq. ft./min. (4 ml./cm²/min.). For contamination of the test solutions <i>E. coli</i> LPS obtained from Sigma Chemical Company was used. Limulus Amebocyte Lysate (LAL) was used for endotoxin determination, and the	5
5	sensitivity for each test solution was found to be 30 pg./ml.  Using the small volume procedure, a number of currently available filters were evaluated for endotoxin removal from 0.9% NaCl. From these results, the filter media of this invention appeared to give the best overall pyrogen removal reducing endotoxin levels to less than the limits of the assay (30 pg./ml.) from small	-
	volumes when the concentration of endotoxin ranged from 1,000 to 1,000,000 pg./ml. The present filters also removed endotoxin from a large variety of solutions of different pH and salt concentrations used for injection. In all cases, the level of endotoxin was reduced below 100 pg./ml. when the original solution was contaminated with 100,000 pg./ml. of endotoxin. The pH of the test solution was found to influence the efficiency of the present filters. Between pH 7.5 and 8.5 in 0.9% NaCl endotoxin removal begins to exceed 100	10
15	pg./ml. However, by using two such filters in series, the level of endotoxin can be reduced to less than 30 pg./ml. in solutions originally contaminated with 1,000,000 pg./ml.  Autoclaving was not found to affect the ability of the present filters to remove endotoxin. The volume of wash solution to remove endotoxin initially present in the filters was found to be from less than 2.6 ml./cm² to no more than 6.4 ml./cm² depending on the nature of the solution.	15
20	In the initial large volume experiment, a comparison of flow rate indicated performance of endotoxin removal was not affected at flow rates of from 3-12 gal./sq. ft./min. when the solution was contaminated with 100,000 pg./ml. At endotoxin concentration of 1,000,000 pg./ml., the present filters were capable of reducing the level of endotoxin below 100 pg./ml. This experiment also indicated that the present filter could remove at least 2.5 mg./cm <sup>2</sup> of endotoxin from 0.9% NaCl pH 7.0. These filters were also found capable of reducing	20
25	the level of endotoxin below 100 pg./ml. from large volumes of distilled water and 20% Dextrose. At high pH, two such filters were used in series to reduce the endotoxin level to below 100 pg./ml.  Changes in solution pH or salt concentration was not found to result in any large scale release of previously adsorbed endotoxin.  In summary, the present filter media are capable of endotoxin removal from heavily contaminated	25
30	solutions used for injection. In all solutions below pH 8.5 these filters reduced endotoxin below 30 pg./ml. when originally contaminated with 100 ng. (100,000 pg./ml.) of endotoxin. Solutions containing 1,000 ng./ml. (1,000,000 pg./ml.) endotoxin can be reduced to below 100 pg./ml. by these filters. With two of these filters in series, endotoxin levels can be kept below 30 pg./ml. in most solutions even with pH's exceeding 8.5. The	30
35	present filter media have greater endotoxin removing ability than comparable filters, e.g., those including cationic melamine-formaldehyde and polyamido-polyamine epichlorodrin as charge modifier on otherwise comparable media.  CLAIMS	35
40	the state of the s	40
45	<ol> <li>The filter media sheet of claim 1, comprising sufficient cationic colloidal silica to provide a positive zeta potential to at least one of said fiber and said particulate.</li> <li>The filter media sheet of claim 1, comprising from about 3 to about 6 percent by weight of said cationic colloidal silica.</li> </ol>	45
50	<ul> <li>5. The filter media sheet of claim 1, wherein said cationic colloidal silica is derived from a dispersion of positively charged colloidal particles having a dense silica core modified with a polyvalent metal-oxygen compound.</li> <li>6. The filter media sheet of claim 5, wherein said polyvalent metal is aluminum, and sufficient of said compound is provided to coat the silica to a level of at least 13% by weight of colloidal solids.</li> </ul>	50
55	7. The filter media sheet of claim 5, wherein the silica dispersion comprises at least 15 percent alumina by weight of colloidal solids.  8. The filter media sheet of claim 1, wherein said fine particulate comprises at least 50 percent by weight of diatomecous earth.	55
60	9. The filter of claim 1, wherein said fine particulate comprises an admixture of diatomaceous earth and perlite.  10. The filter of claim 1, wherein said fine particulate exhibits an average particulate dimension of less than about 10 microns.  11. A method for the preparation of filter media sheets adapted for the removal of electronegative submicron contaminants from contaminated liquids comprising forming a filter sheet from fine particulate	60
65	and a cellulose pulp system as a self-bonding matrix, the surfaces of at least one of said particulate and said pulp being modified with cationic colloidal silica.	65

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Freeness for said system of 100 to 600 ml.

- 13. The method of claim 11, wherein said cationic colloidal silica is derived from a dispersion of positively charged colloidal particles having a dense silica core modified with a polyvalent metal-oxygen compound.
- 14. The method of claim 13, wherein said polyvalent metal is aluminum, and sufficient of said compound is provided to coat the silica to a level of at least 13% by weight of colloidal solids.
- 15. The method of claim 13, wherein the silica dispersion comprises at least 15 percent alumina by weight of colloidal solids.
  - 16. The method of claim 15, further including the step of sanitizing or sterilizing said sheet.
- 10 17. The method of claim 16, wherein the sheet is autoclaved.
  - 18. The method of claim 16, wherein the sheet is hot water flushed.
  - 19. A method for the removal of submicron contaminants from biological fluids comprising passing the contaminated fluid through th filter media sheet of claim 1.
- 20. A method for the removal of submicron contaminants from biological fluids comprising passing the contaminated fluid through a sanitized or sterilized filter sheet comprising particulate material and a self-bonding matrix consisting of a cellulose pulp system, at least one of which is surface modified with a dispersion of cationic colloidal silica comprising at least 15% alumina by weight of the colloidal solids.
  - 21. A filter media sheet substantially as described herein with reference to the accompanying drawings.

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